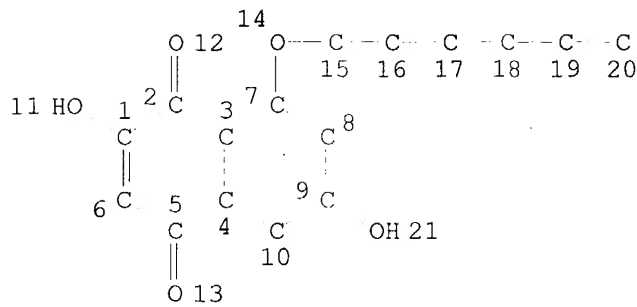


=> d 13 que stat  
L1 STR



NODE ATTRIBUTES:

DEFAULT MLEVEL IS ATOM  
DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED  
NUMBER OF NODES IS 21

STEREO ATTRIBUTES: NONE

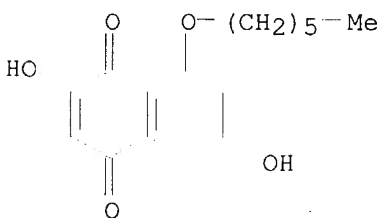
L3 1 SEA FILE=REGISTRY SSS FUL L1

100.0% PROCESSED 40 ITERATIONS  
SEARCH TIME: 00.00.01

1 ANSWERS

=> d ide cbib abs

L3 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2004 ACS on STN  
RN 561297-46-9 REGISTRY  
CN 1,4-Naphthalenedione, 8-(hexyloxy)-2,6-dihydroxy- (9CI) (CA INDEX NAME)  
OTHER NAMES:  
CN Asperaldin  
FS 3D CONCORD  
MF C16 H18 O5  
SR CA  
LC STN Files: CA, CAPLUS, USPATFULL



\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

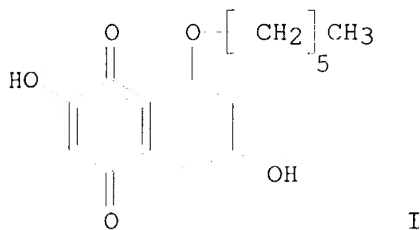
2 REFERENCES IN FILE CA (1907 TO DATE)  
2 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 139:319747 Asperaldin, a new aldose reductase inhibitor from *Aspergillus niger* CFR-1046. I. Fermentation, isolation and

Searched by: Mary Hale 571-272-2507 REM 1D86

characterization. Rao, K. C. Sekhar; Divakar, S.; Srinivas, M.; Babu, K. Naveen; Karanth, N. G.; Sattur, A. P. (Fermentation Technology and Bioengineering Department, Central Food Technological Research Institute, Mysore, 5700013, India). Journal of Antibiotics, 56(2), 173-176 (English) 2003. CODEN: JANTAJ. ISSN: 0021-8820. Publisher: Japan Antibiotics Research Association.

GI

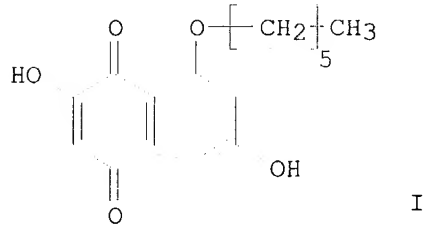


AB The fermentation, isolation, physicochem. properties and biol. activities of asperaldin (I), a new aldose reductase inhibitor, are described. I was produced by *Aspergillus niger* CFR-1046. The EI-MS spectra of the compound showed mol. ions at m/z 205, based on the mass spectra, and giving a mol. formula of C<sub>16</sub>H<sub>18</sub>O<sub>5</sub>, with the chemical name of 2,6-dihydroxy-8-hexyl-oxy-1,4-naphthaquinone. I exhibited a dose-dependent aldose reductase inhibition at an IC<sub>50</sub> of 27  $\mu$ M.

REFERENCE 2: 139:116340 Aldose reductase inhibitor and process for preparation thereof. Sattur, Avinash Prahalad; Rao, Kadiyala Chandrasekhar; Babu, Kilaru Naveen; Soundar, Divakar; Karanth, Naikanakatte Ganesh; Tumkur, Ramachandraiah Shamala (India). U.S. Pat. Appl. Publ. US 2003134399 A1 20030717, 9 pp. (English). CODEN: USXXCO.

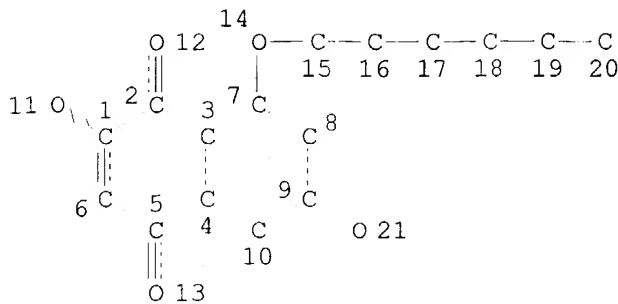
APPLICATION: US 2001-24574 20011221.

GI



AB Aldose reductase inhibitor (I) and pharmaceutically acceptable derivs. thereof derived from cultures of *Aspergillus niger* CFR 1046 and useful as a rat lens aldose reductase inhibitor I are claimed.

=> => d 17 que stat;s 17 not 13  
L5 STR



NODE ATTRIBUTES:

DEFAULT MLEVEL IS ATOM  
DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED  
NUMBER OF NODES IS 21

STEREO ATTRIBUTES: NONE

L7 1 SEA FILE=REGISTRY SSS FUL L5

100.0% PROCESSED 3926 ITERATIONS  
SEARCH TIME: 00.00.01

1 ANSWERS

L8 0 L7 NOT L3

=> e aldose reductase/cn 5

E1 1 ALDOSE MUTAROTASE/CN  
E2 1 ALDOSE OXIDASE/CN  
E3 1 --> ALDOSE REDUCTASE/CN  
E4 1 ALDOSE REDUCTASE (ALFALFA STRAIN RA3/REGEN-S)/CN  
E5 1 ALDOSE REDUCTASE (BARLEY CLONE E3.22-69 GENE AR-H)/CN

=> e

E6 1 ALDOSE REDUCTASE (DIGITALIS PURPUREA GENE AR1)/CN  
E7 1 ALDOSE REDUCTASE (DIGITALIS PURPUREA GENE AR2)/CN  
E8 1 ALDOSE REDUCTASE (EC 1.1.1.21) (ESCHERICHIA COLI O157:H7 STRAIN EDL933 GENE YAFB)/CN  
E9 1 ALDOSE REDUCTASE (HUMAN N-TERMINAL FRAGMENT)/CN  
E10 1 ALDOSE REDUCTASE (MOUSE REDUCED)/CN  
E11 1 ALDOSE REDUCTASE (MOUSE RENAL-SPECIFIC)/CN  
E12 1 ALDOSE REDUCTASE (MOUSE STRAIN SV129J CLONE MAR-F GENE ALDOR 1)/CN  
E13 1 ALDOSE REDUCTASE (MUS MUSCULUS CLONE KE2)/CN  
E14 1 ALDOSE REDUCTASE (PIG LENS) (EC 1.1.1.21)/CN  
E15 1 ALDOSE REDUCTASE (RAT RENAL-SPECIFIC)/CN  
E16 1 ALDOSE REDUCTASE (RENAL-SPECIFIC HUMAN)/CN  
E17 1 ALDOSE REDUCTASE (RHODOPSEUDOMONAS PALUSTRIS CGA009 STRAIN CGA009 GENE YAFB)/CN

=> s aldose reductase?/cn

L9 23 ALDOSE REDUCTASE?/CN

=> e cfr 1046/cn 5

E1 1 CFPR-G 200/CN

Searched by: Mary Hale 571-272-2507 REM 1D86

E2 1 CFPRBK 708S/CN  
E3 0 --> CFR 1046/CN  
E4 1 CFR 2/CN  
E5 1 CFR 20/30/CN

=> s dihydroxy(1)hexoxy(1)naphthaquinone  
323484 DIHYDROXY  
191 HEXOXY  
16 NAPHTHAQUINONE  
L10 0 DIHYDROXY(L)HEXOXY(L)NAPHTHAQUINONE

=> aspergillus niger/cn 5  
ASPERGILLUS IS NOT A RECOGNIZED COMMAND  
The previous command name entered was not recognized by the system.  
For a list of commands available to you in the current file, enter  
"HELP COMMANDS" at an arrow prompt (=>).

=> e aspergillus niger/cn 5  
E1 1 ASPERGILLUS MELLEUS SEMI-ALKALINE PROTEINASE/CN  
E2 1 ASPERGILLUS NIDULANS NEUTRAL PROTEINASE/CN  
E3 1 --> ASPERGILLUS NIGER/CN  
E4 1 ASPERGILLUS NIGER ACID PROTEASE/CN  
E5 1 ASPERGILLUS NIGER ACID PROTEINASE/CN

=> s e3  
L11 1 "ASPERGILLUS NIGER"/CN

=> fil medl,hcapl,biosis,embase;s (19 or aldose reductase?) (1)(l11 or aspergill?  
niger or cfr 1046)

COST IN U.S. DOLLARS	SINCE FILE ENTRY	TOTAL SESSION
FULL ESTIMATED COST	339.70	340.12
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE ENTRY	TOTAL SESSION
CA SUBSCRIBER PRICE	-0.66	-0.66

FILE 'MEDLINE' ENTERED AT 09:21:02 ON 19 MAR 2004

FILE 'HCAPLUS' ENTERED AT 09:21:02 ON 19 MAR 2004  
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L12 2 FILE MEDLINE  
L13 7 FILE HCAPLUS  
L14 2 FILE BIOSIS  
L15 2 FILE EMBASE

TOTAL FOR ALL FILES  
L16 13 (L9 OR ALDOSE REDUCTASE?) (L) (L11 OR ASPERGILL? NIGER OR CFR  
1046)

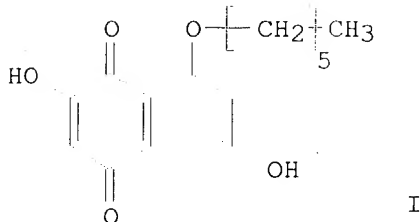
=> dup rem 116  
PROCESSING COMPLETED FOR L16

L17 7 DUP REM L16 (6 DUPLICATES REMOVED)

=> d 1-7 cbib abs

L17 ANSWER 1 OF 7 HCPLUS COPYRIGHT 2004 ACS on STN  
2003:551102 Document No. 139:116340 Aldose reductase inhibitor and process  
for preparation thereof. Sattur, Avinash Prahalad; Rao, Kadiyala  
Chandrasekhar; Babu, Kilaru Naveen; Soundar, Divakar; Karanth,  
Naikanakatte Ganesh; Tumkur, Ramachandraiah Shamala (India). U.S. Pat.  
Appl. Publ. US 2003134399 A1 20030717, 9 pp. (English). CODEN: USXXCO.  
APPLICATION: US 2001-24574 20011221.

GI



AB Aldose reductase inhibitor (I) and pharmaceutically acceptable derivs. thereof derived from cultures of **Aspergillus niger** CFR 1046 and useful as a rat lens aldose reductase inhibitor I are claimed.

L17 ANSWER 2 OF 7 HCPLUS COPYRIGHT 2004 ACS on STN  
2003:420474 Document No. 139:226997 Isolation and characterization of two specific regulatory *Aspergillus niger* mutants shows antagonistic regulation of arabinan and xylan metabolism. de Groot, Marco J. L.; van de Vondervoort, Peter J. I.; de Vries, Ronald P.; van Kuyk, Patricia A.; Ruijter, George J. G.; Visser, Jaap (Section Molecular Genetics of Industrial Micro-organisms, Wageningen University, Wageningen, NL-6703HA, Neth.). Microbiology (Reading, United Kingdom), 149(5), 1183-1191 (English) 2003. CODEN: MROBEO. ISSN: 1350-0872. Publisher: Society for General Microbiology.

AB This paper describes two *Aspergillus niger* mutants (araA and araB) specifically disturbed in the regulation of the arabinanase system in response to the presence of L-arabinose. Expression of the three known L-arabinose-induced arabinanolytic genes, abfA, abfB and abnA, was substantially decreased or absent in the araA and araB strains compared to the wild-type when incubated in the presence of L-arabinose or L-arabitol. In addition, the intracellular activities of L-arabitol dehydrogenase and L-arabinose reductase, involved in L-arabinose catabolism, were decreased in the araA and araB strains. Finally, the data show that the gene encoding D-xylulose kinase, xkiA, is also under control of the arabinanolytic regulatory system. L-Arabitol, most likely the true inducer of the arabinanolytic and L-arabinose catabolic genes, accumulated to a high intracellular concentration in the araA and araB mutants. This indicates that the decrease of expression of the arabinanolytic genes was not due to lack of inducer accumulation. Therefore, it is proposed that the araA and araB mutations are localized in pos.-acting components of the regulatory system involved in the expression of the arabinanase-encoding genes and the genes encoding the L-arabinose catabolic pathway.

L17 ANSWER 3 OF 7 MEDLINE on STN DUPLICATE 1  
2003196168. PubMed ID: 12715878. Asperaldin, a new aldose reductase inhibitor from *Aspergillus niger* CFR-

1046. I. Fermentation, isolation and characterization. Rao K C  
Sekhar; Divakar S; Srinivas M; Babu K Naveen; Karanth N G; Sattur A P.  
(Fermentation Technology and Bioengineering Department, Central Food  
Technological Research Institute, Mysore 5700013, India. ) Journal of  
antibiotics, (2003 Feb) 56 (2) 173-6. Journal code: 0151115. ISSN:  
0021-8820. Pub. country: Japan. Language: English.

L17 ANSWER 4 OF 7 MEDLINE on STN DUPLICATE 2  
2002697455. PubMed ID: 12458767. Nigerloxin, a novel inhibitor of  
**aldose reductase** and lipoxygenase with Free radical  
scavenging activity from **Aspergillus niger** CFR-W-105.  
Rao K C Sekhar; Divakar S; Babu K Naveen; Rao A G Appu; Karanth N G;  
Sattur A P. (Fermentation Technology and Bioengineering Department,  
Central Food Technological Research Institute, Mysore 570 013, India. )  
Journal of antibiotics, (2002 Sep) 55 (9) 789-93. Journal code: 0151115.  
ISSN: 0021-8820. Pub. country: Japan. Language: English.

AB An enzyme inhibitor, nigerloxin, with inhibition against soy bean  
lipoxygenase-I (LOX-1), rat lens **aldose reductase**  
(RLAR) as well as free radical scavenging activity was isolated from the  
fermented wheat bran using **Aspergillus niger**  
CFR-W-105. Its chemical structure was identified as 2-amido-3-hydroxy-6-  
methoxy-5-methyl-4-(prop-1'-enyl) benzoic acid by NMR and GCEIMS data.  
The IC<sub>50</sub> values against LOX-1 and RLAR were found to be 79 microM and 69  
microM and ED<sub>50</sub> against 1,1-diphenyl-2-picrylhydrazyl (DPPH) was 66  
microM.

L17 ANSWER 5 OF 7 HCPLUS COPYRIGHT 2004 ACS on STN  
1997:628724 Document No. 127:316693 Isolation of *Aspergillus niger* creA  
mutants and effects of the mutations on expression of arabinases and  
L-arabinose catabolic enzymes. Ruijter, George J. G.; Vanhanen, Sipo A.;  
Gielkens, Marco M. C.; van de Vondervoort, Peter J. I.; Visser, Jaap  
(Section Molecular Genetics of Industrial Microorganisms, Wageningen  
Agricultural University, Wageningen, 6703 HA, Neth.). Microbiology  
(Reading, United Kingdom), 143(9), 2991-2998 (English) 1997. CODEN:  
MROBEO. ISSN: 1350-0872. Publisher: Society for General Microbiology.

AB *Aspergillus niger* mutants relieved of carbon repression were isolated from  
an areA parental strain by selection of colonies that exhibited improved  
growth on a combination of 4-aminobutanoic acid (GABA) and D-glucose. In  
addition to derepression of the utilization of GABA as a nitrogen source in  
the presence of D-glucose, three of the four mutants also showed  
derepression of L-alanine and L-proline utilization. Transformation of  
the mutants with the *A. niger* creA gene, encoding the repressor protein  
CREA, re-established the areA phenotype on GABA/D-glucose, identifying the  
mutations as creAd. The creA gene mapped on chromosome IV by linkage  
anal. and contour-clamped homogeneous elec. field hybridization. The creA  
mutants obtained were used to study the involvement of CREA in repression  
by D-glucose of arabinases and L-arabinose catabolism in *A. niger*. In  
wild-type *A. niger*,  $\alpha$ -L-arabinofuranosidase A,  $\alpha$ -L-  
arabinofuranosidase B, endo-arabinase, L-arabinose reductase and  
L-arabitol dehydrogenase were induced on L-arabinose, but addition of  
D-glucose prevented this induction. Repression was relieved to varying  
degrees in the creA mutants, showing that biosynthesis of arabinases and  
L-arabinose catabolic enzymes is under control of CREA.

L17 ANSWER 6 OF 7 HCPLUS COPYRIGHT 2004 ACS on STN  
1993:187567 Document No. 118:187567 Induction of extracellular arabinases on  
monomeric substrates in *Aspergillus niger*. Van der Veen, Peter; Flippes,  
Michel J. A.; Voragen, Alphons G. J.; Visser, Jaap (Dep. Genet., Agric.  
Univ., Wageningen, 6703 HA, Neth.). Archives of Microbiology, 159(1),  
66-71 (English) 1993. CODEN: AMICCW. ISSN: 0302-8933.

AB The induction of extracellular arabinases by pentose sugars and polyols  
generated by the metabolic pathway of L-arabinose and D-xylose catabolism

in *Aspergillus niger* was investigated. Induction occurred with L-arabinose and L-arabitol but not with D-xylose or xylitol. L-Arabitol, in particular, was found to be a good inducer for  $\alpha$ -L-arabinofuranosidase and endo-arabinose activities. Western blotting anal. showed both  $\alpha$ -L-arabinofuranosidase A and B to be present. No induction was observed using D-arabitol. Unlike the wild-type *A. niger* N402 strain, the *A. niger* xylulose kinase-neg. mutant N572 also showed induction of  $\alpha$ -L-arabinofuranosidases A and B and endo-arabinose activity on D-xylose and xylitol. This is due to metabolic conversion of these compds. leading to the accumulation of both xylitol and L-arabitol in this mutant, the latter of which then acts as inducer. The induction of the two  $\alpha$ -L-arabinofuranosidases and endo-arabinase is under the control of two regulatory systems, namely pathway specific induction and carbon catabolite repression. Under derepressing conditions in the wild type, only a  $\alpha$ -L-arabinofuranosidase B could be detected by Western blotting anal. This indicates that  $\alpha$ -L-arabinofuranosidase B is of importance in the initiation of specific induction of the various arabinose activities in *A. niger* grown on arabinose-containing structural polysaccharides.

L17 ANSWER 7 OF 7 HCAPLUS COPYRIGHT 2004 ACS on STN

1989:591175 Document No. 111:191175 L-Arabinose and D-xylose catabolism in *Aspergillus niger*. Witteveen, C. F. B.; Busink, R.; Van de Vonderboort, P.; Dijkema, C.; Swart, K.; Visser, J. (Dep. Genet., Agric. Univ., Wageningen, 6703 HA, Neth.). Journal of General Microbiology, 135(8), 2163-71 (English) 1989. CODEN: JGMIAN. ISSN: 0022-1287.

AB A mutant of *A. niger* unable to grow on D-xylose and L-arabinose was isolated. Genetic anal. revealed that the mutation is located on linkage group IV. Enzymic anal. revealed a deficiency in D-xylulose kinase activity. After transfer of growing mycelium to a medium containing either D-xylose or L-arabinose, the mutant accumulates large amts. of arabitol and xylitol, as shown by  $^{13}\text{C}$  NMR spectroscopy. These data and an anal. of enzyme activities induced by D-xylose and L-arabinose in the wild-type strain led to the following catabolic pathway for D-xylose: D-xylose-xylitol-D-xylulose-D-xylulose 5-phosphate; and for L-arabinose: L-arabinose-L-arabitol-L-xylulose-xylitol-D-xylulose-D-xylulose 5-phosphate. The reduction steps of the sugars to the corresponding polyols are all NADPH dependent. The oxidation steps of the polyols to the sugars are all NAD<sup>+</sup> dependent. Fractionation of cell-free exts. gave information about the specificity of the enzymes and showed that all the reactions are catalyzed by different enzymes.

=> s sattur, a?/au;s rao, k?/au;s babu, k?/au

L18 4 FILE MEDLINE

L19 21 FILE HCAPLUS

L20 16 FILE BIOSIS

L21 7 FILE EMBASE

TOTAL FOR ALL FILES

L22 48 SATTUR, A?/AU

L23 1999 FILE MEDLINE

L24 7052 FILE HCAPLUS

L25 3691 FILE BIOSIS

L26 1419 FILE EMBASE

TOTAL FOR ALL FILES

L27 14161 RAO, K?/AU

L28. 149 FILE MEDLINE  
L29 445 FILE HCAPLUS  
L30 235 FILE BIOSIS  
L31 92 FILE EMBASE

TOTAL FOR ALL FILES  
L32 921 BABU, K?/AU

=> s 122 and 127 and 132  
L33 2 FILE MEDLINE  
L34 3 FILE HCAPLUS  
L35 2 FILE BIOSIS  
L36 1 FILE EMBASE

TOTAL FOR ALL FILES  
L37 8 L22 AND L27 AND L32

=> s 137 not 116  
L38 0 FILE MEDLINE  
L39 0 FILE HCAPLUS  
L40 0 FILE BIOSIS  
L41 0 FILE EMBASE

TOTAL FOR ALL FILES  
L42 0 L37 NOT L16

COST IN U.S. DOLLARS	SINCE FILE ENTRY	TOTAL SESSION
FULL ESTIMATED COST	19.34	359.46
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE ENTRY	TOTAL SESSION
CA SUBSCRIBER PRICE	-3.47	-4.13

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DICTIONARY FILE UPDATES: 17 MAR 2004 HIGHEST RN 664302-53-8

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 6, 2004

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Experimental and calculated property data are now available. For more  
information enter HELP PROP at an arrow prompt in the file or refer  
to the file summary sheet on the web at:  
<http://www.cas.org/ONLINE/DBSS/registryss.html>

=> fil hcap;s "ec 1.1.1.9" or "ec 1.1" or "ec 1.3" or "ec 1.2" or 9028-16-4/rn  
COST IN U.S. DOLLARS SINCE FILE TOTAL  
FULL ESTIMATED COST ENTRY SESSION  
2.10 361.56

Searched by: Mary Hale 571-272-2507 REM 1D86

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE ENTRY	TOTAL SESSION
CA SUBSCRIBER PRICE	0.00	-4.13

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FILE COVERS 1907 - 19 Mar 2004 VOL 140 ISS 13  
 FILE LAST UPDATED: 18 Mar 2004 (20040318/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

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2927 "ECS"
80207 "EC"
  ("EC" OR "ECS")
7837158 "1"
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1667263 "9"
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2927 "ECS"
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7837158 "1"
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L43 8045 "EC 1.1.1.9" OR "EC 1.1" OR "EC 1.3" OR "EC 1.2" OR 9028-16-4/RN

=> s l43 and (inhibit? or modulat?)  
1640208 INHIBIT?  
282129 MODULAT?  
L44 2554 L43 AND (INHIBIT? OR MODULAT?)

=> s l44 py=>2001  
MISSING OPERATOR L44 PY=>2001  
The search profile that was entered contains terms or  
nested terms that are not separated by a logical operator.

=> s l44 and py=>2001  
3208724 PY=>2001  
L45 113 L44 AND PY=>2001

=> d

L45 ANSWER 1 OF 113 HCAPLUS COPYRIGHT 2004 ACS on STN  
AN 2004:175032 HCAPLUS  
TI Are NADP-dependent isocitrate dehydrogenases and ferredoxin-dependent  
glutamate synthase co-regulated by the same photoreceptors?  
AU Appenroth, Klaus-J.; Teller, Steffen  
CS Institute of General Botany and Plant Physiology, University of Jena,  
Dornburger Str. 159, Jena, 07743, Germany  
SO Planta (2004), 218(5), 775-783  
CODEN: PLANAB; ISSN: 0032-0935  
PB Springer-Verlag  
DT Journal  
LA English

=> s l44 and =<2001  
MISSING TERM 'AND =<2001'  
The search profile that was entered contains a logical  
operator followed immediately by another operator.

=> s l44 and py=<2001  
21547848 PY=<2001  
L46 2475 L44 AND PY=<2001

=> d

L46 ANSWER 1 OF 2475 HCAPLUS COPYRIGHT 2004 ACS on STN  
AN 2003:383727 HCAPLUS  
DN 139:191102  
TI Effects of volatile oil constituents of Nigella sativa on carbon  
tetrachloride-induced hepatotoxicity in mice: Evidence for antioxidant  
effects of thymoquinone  
AU Mansour, M. A.; Ginawi, O. T.; El-Hadiyah, T.; El-Khatib, A. S.;  
Al-Shabanah, O. A.; Al-Sawaf, H. A.  
CS Department of Pharmacology, College of Pharmacy, King Saud University,  
Riyadh, 11451, Saudi Arabia  
SO Research Communications in Molecular Pathology and Pharmacology (2001), 110(3 & 4), 239-251  
CODEN: RCMPE6; ISSN: 1078-0297  
PB PJD Publications Ltd.

DT Journal

LA English

RE.CNT 50 THERE ARE 50 CITED REFERENCES AVAILABLE FOR THIS RECORD  
ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> d 2475

L46 ANSWER 2475 OF 2475 HCPLUS COPYRIGHT 2004 ACS on STN  
AN 1962:55214 HCPLUS  
DN 56:55214  
OREF 56:10554e-i  
TI Ribitol dehydrogenase. III. Kinetic studies with product inhibition  
AU Fromm, Herbert J.; Nelson, Dennis R.  
CS Univ. of North Dakota, Grand Forks  
SO Journal of Biological Chemistry (1962), 237, 215-20  
CODEN: JBCHA3; ISSN: 0021-9258  
DT Journal  
LA Unavailable

=> s 146 and addition? product?

MISSING TERM AFTER PRODUCT?  
Operators must be followed by a search term, L-number, or query name.

=> s 146 and addition? product?

161458 ADDITION?  
1375200 ADDN  
68367 ADDNS  
1418632 ADDN  
(ADDN OR ADDNS)  
417064 ADDNL  
1 ADDNLS  
417065 ADDNL  
(ADDNL OR ADDNLS)

1873257 ADDITION?  
(ADDITION? OR ADDN OR ADDNL)

2471606 PRODUCT?

14004 ADDITION? PRODUCT?  
(ADDITION? (W) PRODUCT?)

L47 2 L46 AND ADDITION? PRODUCT?

=> d 1-2 cbib abs

L47 ANSWER 1 OF 2 HCPLUS COPYRIGHT 2004 ACS on STN

1982:2847 Document No. 96:2847 Pig brain aldose reductase: a kinetic study using the centrifugal fast analyzer. Boghosian, Robert A.; McGuinness, Eugene T. (Dep. Chem., Seton Hall Univ., South Orange, NJ, 07079, USA). International Journal of Biochemistry, 13(8), 909-14 (English)

1981. CODEN: IJBOBV. ISSN: 0020-711X.

AB Initial velocity and product inhibition studies of pig brain aldose reductase (EC 1.1.1.21) previously purified to apparent homogeneity, using D-xylose as substrate, indicated a sequential mechanism, probably with an ordered bi bi or an iso Theorell-Chance pattern of substrate addition-product release. The Km values for xylose and NADPH were 4.1 mM and 3.1  $\mu$ M, resp. The advantages of using the centrifugal fast analyzer for reaction rate studies with enzymes, e.g. simultaneous multiple-reaction initiation and parallel monitoring, are discussed.

L47 ANSWER 2 OF 2 HCPLUS COPYRIGHT 2004 ACS on STN

Searched by: Mary Hale 571-272-2507 REM 1D86

1979:485923 Document No. 91:85923 The reaction of carbonyl cyanide phenylhydrazone with thiols. Drobnička, L.; Sturdík, E. (Dep. Microbiol. Biochem., Slovak Polytech. Univ., Bratislava, 880 37, Czech.). Biochimica et Biophysica Acta, 585(3), 462-76 (English) 1979. CODEN: BBACAQ. ISSN: 0006-3002.

AB Carbonyl cyanide phenylhydrazone and its ring-substituted analogs reacted with thiols (thioglycolic acid, 2-mercaptoethanol, dithiothreitol) and aminothiols (cysteine, glutathione) to give the corresponding N-(substituted phenyl)-N'-(alkylthiodicyano)-methylhydrazine derivs. These **addition products** decomposed to the original components in alkaline solution. In the presence of excess thiol in aqueous buffered systems, the addition reactions are practically quant. with respect to phenylhydrazone, follow pseudo-1st-order kinetics, and can be investigated spectrophotometrically. These reactions are of the bimol. AdN type where the nondissocd. forms of carbonyl cyanide phenylhydrazone function as electrophilic components and the RS- ion is the nucleophilic component (attack of the azomethine group). The reactivity of carbonyl cyanide phenylhydrazone with respect to thiols increases in the order carbonyl cyanide phenylhydrazone < carbonyl cyanide m-chlorophenylhydrazone < carbonyl cyanide p-trifluoromethoxyphenylhydrazone, which corresponds to the decreasing order of their pKa values. On the other hand, the reactivity of the thiols increases with their basicity. The reactivity of carbonyl cyanide phenylhydrazone with thiols is comparable to the reactivity of Ph isothiocyanate and N-ethylmaleimide. Carbonyl cyanide phenylhydrazone was an efficient **inhibitor** of rabbit muscle glyceraldehyde 3-phosphate dehydrogenase (EC 1.2.1.12). The results are discussed in relation to the biol. activity of carbonyl cyanide phenylhydrazone.

=> log y			
COST IN U.S. DOLLARS	SINCE FILE	TOTAL	
	ENTRY	SESSION	
FULL ESTIMATED COST	24.78	386.34	
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL	
CA SUBSCRIBER PRICE	ENTRY	SESSION	
	-1.39	-5.52	

STN INTERNATIONAL LOGOFF AT 09:31:02 ON 19 MAR 2004